## AMENDMENTS TO THE CLAIMS

## IN THE CLAIMS:

The following claim listing is meant to replace all previous claim listings.

- 1. (Withdrawn) A method for the design and/or the selection of chemokines variants having agonist or antagonist property towards a ligandof GPCR of animal cells comprising the following steps:
  - A) obtaining a phage displayed library expressing on their surface said chemokine variants mutated within the domain responsible for their effector function,
  - B) having a culture of animal cells expressing on their membranes the GPCR,
  - C) Incubating the cell culture with the phage library obtained In A),
  - D) harvesting the cells after removal of non specifically bond and surface receptor bound phages,
  - E) Releasing the phages internalized in step C) by lysis of cells obtained in D)
  - F) Infecting an *E. coli* culture with the released phages obtained in E) and amplifying the clones previously internalized,
  - G) Obtaining a phage library enriched in internalizing chemokines ligands,
  - H) Assaying the agonist or antagonist property of the chemokine variants versus the native one.
- 2. (Withdrawn) The method according to claim 1 wherein the chemokine is RANTES.
- 3. (Withdrawn) The method according to claim 1 wherein the GPCR expressed within the membrane of animal cells is CCR5.

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4. (Withdrawn) The method according to claim 1 wherein the animal cells are human cells.

- 5. (Withdrawn) The method according to claim 2 wherein the phage library of RANTES variants is obtained using a method comprising the following steps:
  - Obtaining a DNA sequence coding for human RANTES resulting from the ampification of cDNA prepared from activated PBMCs,
  - Performing a PCR mutagenesis of the 5' portion of the DNA sequence of RANTES using a specific downstream primer and a degenerate upstream primer containing recognition sites for restriction enzymes in order to insert the PCR amplification products into the phage display vector,
  - Inserting the purified PCR products into a phage display vector,
  - Production of the phage library by introducing the vector containing the purified PCR products into an *E. coli* culture.
  - 6. (Withdrawn) The method according to claim 2 wherein anti-HIV activity is assayed.
- 7. (Withdrawn) A method for the design and/or the selection of chemokines having agonist or antagonist property towards a GPCR of animal cells comprising the following steps:
  - A. obtaining a phage displayed library expressing on their surface said chemokine mutated within the domain responsible for their effector function
  - B. having a culture of animal cells expressing on their membranes the GPCR,
  - C. Incubating the cell culture with the phage library obtained in A),
  - D. Eliminating the non specifically bond phages from the cells, by a process keeping the specifically bound phages on the said receptor

E. Incubating the cells obtained in D) with an E. coli culture and amplifying the clones being infected by the phages bound to the said receptor on animal cells,

- F. Obtaining a phage library enriched in externally bound phages,
- G. Assaying the agonist or antagonist property of the chemokine variants versus the native chemokine.
- 8. (Withdrawn) The method according to claim 7, wherein the chemokine is RANTES.
- 9. (Withdrawn) The method according to claim 7, wherein the GPCR expressed within the membrane of animal cells is CCR5.
- 10. (Withdrawn) The method according to claim 7, wherein the animal cells are human cells.
- 11. (Withdrawn)The method according to claim 8, wherein the phage library of RANTES variants is obtained using a method comprising the following steps:
  - Obtaining a DNA sequence coding for human RANTES resulting from the ampification of cDNA prepared from activated PBMCs,
  - Performing a PCR mutagenesis of the 5'portion of the DNA sequence of RANTES
    using a specific downstream primer and a degenerate upstream primer containing
    recognition sites for restriction enzymes in order to insert the PCR amplification
    products into the phage display vector,

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- Inserting the purified PCR products into a phage display vector,

- Production of the phage library by introducing the vector containing the purified PCR products into an E; coli culture.
- 12. (Withdrawn) The method according to claim 8 wherein anti-HIV activity is assayed.
- 13. (Currently Amended) A compound comprising the following formula: XaaSPXaa Xaa, Xaa, Xaa (SEQ ID NO:40) Xaa Ser Pro Xaa Ser Ser Gln Xaa Xaa Xaa RANTES 10-68 (SEQ ID NO: 41) in which
  - Xaa at position 1 is L or an aromatic residue,
  - Xaa at position 4 is L, M or V
  - Xaa at position 8-10 is S, P, T or A.
- 14. (Previously Presented) The compound according to claim 13 having one of the following formulae :

```
LSPVSSQSSA (SEQ ID NO: 1) (P<sub>1</sub>)
FSPLSSQSSA (SEQ ID NO: 2) (P<sub>2</sub>)
LSPMSSQSPA (SEQ ID NO: 3)
WSPLSSQSPA (SEQ ID NO: 4)
WSPLSSQSSP (SEQ ID NO: 5)
LSPLSSQSAA (SEQ ID NO: 15)
YSPLSSQSSP (SEQ ID NO: 17)
```

15. (Withdrawn) The compound according to claim 13 having the formula: FSPLSSQSSA(SEQ ID N): 2-RANTES(10-68).

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16.(Withdrawn) The compound according to claim 13 having the formula: LSPVSSQSSA-RANTES (10-68).

17.(Currently Amended) A pharmaceutical composition which comprises of a compound having the formula: XaaSPXaa Xaa, Xaa, Xaa (SEQ ID NO:40) Xaa Ser Pro Xaa Ser Gln Xaa Xaa Xaa - RANTES 10-68 (SEQ ID NO: 41) in which

- Xaa at position 1 is L or an aromatic residue,
- Xaa at position 4 is L, M or V
- Xaa at position 8-10 is S, P, T or A.

or a pharmaceutical salt thereof, in a mixture with one or more pharmaceutically acceptable excipient.

18.(Withdrawn) The composition of claim 17, in which the compound have the formula: LSPVSSQSSA(SEQ ID NO: 1)- RANTES(10-68).

19.(Withdrawn) The composition of claim 17, in which the compound have the formula: FSPLSSQSSA (SEQ ID NO:2) -RANTES-(10-68).

20.(Withdrawn) A method for preventing and/or inhibiting HIV infection in humans comprising a step of treatment with a composition of claim 18.

21.(Withdrawn) A method for preventing and/or inhibiting HIV infection in humans comprising a step of treatment with a composition of claim 19.

22.(Withdrawn)A method for preventing and/or curing inflammatory or malignant diseases in humans comprising a step of treatment with a composition of claim 13 or 14.

23. (Previously Presented) A compound comprising one of the following formulae:

```
LSPQSSLSSS
                  (SEQ ID NO: 6),
                  (SEQ ID NO: 7),
ASSGSSQSTS
ISAGSSQSTS
                  (SEQ ID NO: 8),
RSPMSSQSSP
                  (SEQ ID NO: 9),
                  (SEQ ID NO: 10),
YSPSSSLAPA
                  (SEQ ID NO: 11),
MSPLSSQASA
ASPMSSQSSS
                  (SEQ ID NO: 12),
                  (SEQ ID NO: 13),
QSPLSSQAST
                  (SEQ ID NO: 14),
QSPLSSTASS
                  (SEQ ID NO: 16),
GSSSSSQTPA
FSSVSSQSSS,
                  (SEQ ID NO: 18),
                  (SEQ ID NO: 30),
VSTLSSPAST,
                  (SEQ ID NO: 31),
ASSFSSRAPP,
                  (SEQ ID NO: 32),
QSSASSSSSA
                  (SEQ ID NO: 33),
QSPGSSWSAA,
                  (SEQ ID NO: 34),
QSPPSSWSSS,
                  (SEQ ID NO: 35) and
QSPLSSFTSS,
                  (SEQ ID NO: 36).
ASPQSSLPAA,
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24. (Previously Presented) A compound consisting essentially of one of the following formulae:

```
LSPQSSLSSS
                  (SEQ ID NO: 6),
                  (SEQ ID NO: 7),
ASSGSSQSTS
                  (SEQ ID NO: 8),
ISAGSSQSTS
                  (SEQ ID NO: 9),
RSPMSSQSSP
YSPSSSLAPA
                  (SEQ ID NO: 10),
                  (SEQ ID NO: 11),
MSPLSSQASA
                  (SEQ ID NO: 12),
ASPMSSQSSS
QSPLSSQAST
                  (SEQ ID NO: 13),
QSPLSSTASS
                  (SEQ ID NO: 14),
                  (SEQ ID NO: 16),
GSSSSSQTPA
FSSVSSQSSS,
                  (SEQ ID NO: 18),
VSTLSSPAST,
                  (SEQ ID NO: 30),
ASSFSSRAPP,
                 (SEQ ID NO: 31),
                 (SEQ ID NO: 32),
QSSASSSSSA
QSPGSSWSAA,
                 (SEQ ID NO: 33),
                 (SEQ ID NO: 34),
QSPPSSWSSS,
                 (SEQ ID NO: 35) and
QSPLSSFTSS,
                 (SEQ ID NO: 36).
ASPQSSLPAA,
```

25. (Previously Presented) A pharmaceutical composition which comprises one of the following formulae:

```
LSPQSSLSSS (SEQ ID NO: 6),
ASSGSSQSTS (SEQ ID NO: 7),
ISAGSSQSTS (SEQ ID NO: 8),
RSPMSSQSSP (SEQ ID NO: 9),
YSPSSSLAPA (SEQ ID NO: 10),
```

```
(SEQ ID NO: 11),
MSPLSSQASA
                  (SEQ ID NO: 12),
ASPMSSQSSS
                  (SEQ ID NO: 13),
QSPLSSQAST
QSPLSSTASS
                  (SEQ ID NO: 14),
                  (SEQ ID NO: 16),
GSSSSSQTPA
                  (SEQ ID NO: 18),
FSSVSSQSSS,
VSTLSSPAST,
                  (SEQ ID NO: 30),
                  (SEQ ID NO: 31),
ASSFSSRAPP,
                  (SEQ ID NO: 32),
QSSASSSSSA
                  (SEQ ID NO: 33),
QSPGSSWSAA,
                  (SEQ ID NO: 34),
QSPPSSWSSS,
                  (SEQ ID NO: 35) and
QSPLSSFTSS,
                  (SEQ ID NO: 36).
ASPQSSLPAA,
```

or a pharmaceutical salt thereof, in a mixture with one or more pharmaceutically acceptable excipient.

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26. (Previously Presented) A pharmaceutical composition consisting essentially of one of the following formulae:

```
(SEQ ID NO: 6),
LSPQSSLSSS
                  (SEQ ID NO: 7),
ASSGSSQSTS
                  (SEQ ID NO: 8),
ISAGSSQSTS
                  (SEQ ID NO: 9),
RSPMSSQSSP
                  (SEQ ID NO: 10),
YSPSSSLAPA
                  (SEQ ID NO: 11),
MSPLSSQASA
                  (SEQ ID NO: 12),
ASPMSSQSSS
                  (SEQ ID NO: 13),
QSPLSSQAST
QSPLSSTASS
                  (SEQ ID NO: 14),
                  (SEQ ID NO: 16),
GSSSSSQTPA
                  (SEQ ID NO: 18),
FSSVSSQSSS,
```

VSTLSSPAST,	(SEQ ID NO: 30),
ASSFSSRAPP,	(SEQ ID NO: 31),
QSSASSSSSA	(SEQ ID NO: 32),
QSPGSSWSAA	(SEQ ID NO: 33),
QSPPSSWSSS,	(SEQ ID NO: 34),
QSPLSSFTSS,	(SEQ ID NO: 35) and
ASPQSSLPAA,	(SEQ ID NO: 36).

or a pharmaceutical salt thereof, in a mixture with one or more pharmaceutically acceptable excipient.